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An integrated, directed mass spectrometric approach for in-depth characterization of complex peptide mixtures

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Abstract: LC-MS/MS has emerged as the method of choice for the identification and quantification of protein sample mixtures. For very complex samples such as complete proteomes, the most commonly used LC-MS/MS method, data-dependent acquisition (DDA) precursor selection, is of limited utility. The limited scan speed of current mass spectrometers along with the highly redundant selection of the most intense precursor ions generates a bias in the pool of identified proteins toward those of higher abundance. A directed LC-MS/MS approach that alleviates the limitations of DDA precursor ion selection by decoupling peak detection and sequencing of selected precursor ions is presented. In the first stage of the strategy, all detectable peptide ion signals are extracted from high resolution LC-MS feature maps or aligned sets of feature maps. The selected features or a subset thereof are subsequently sequenced in sequential, non-redundant directed LC-MS/MS experiments, and the MS/MS data are mapped back to the original LC-MS feature map in a fully automated manner. The strategy, implemented on an LTQ-FT MS platform, allowed the specific sequencing of 2,000 features per analysis and enabled the identification of more than 1,600 phosphorylation sites using a single reversed phase separation dimension without the need for time-consuming prefractionation steps. Compared with conventional DDA LC-MS/MS experiments, a substantially higher number of peptides could be identified from a sample, and this increase was more pronounced for low intensity precursor ions.

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An integrated, directed mass spectrometric approach for in-depth
characterization of complex peptide mixtures

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Abbreviations:

DDA: data dependent acquisition, PBS: phosphate buffered saline, TCEP: tris(2-carboxyethyl)phosphine, FA: formic acid, MPS: monoisotopic precursor selection, INL: inclusion list, NL: neutral loss

Summary

Mass spectrometry in combination with liquid chromatography (LC-MS/MS) has emerged as the method of choice for the identification and quantification of protein sample mixtures. For very complex samples such as complete proteomes, the most commonly used LC-MS/MS method, data dependent (DDA) precursor selection, is of limited utility. The limited scan speed of current mass spectrometers, along with the highly redundant selection of the most intense precursor ions generate a bias in the pool of identified proteins towards those of higher abundance. A directed LC-MS/MS approach that alleviates the limitations of DDA precursor ion selection by decoupling peak detection and sequencing of selected precursor ions is presented. In the first stage of the strategy, all detectable peptide ion signals are extracted from high resolution LC-MS feature maps or aligned sets of feature maps. The selected features or a subset thereof are subsequently sequenced in sequential, non-redundant directed LC-MS/MS experiments and the MS/MS data are mapped back to the original LC-MS feature map in a fully automated manner. The strategy, implemented on a LTQ-FT MS platform, allowed the specific sequencing of 2,000 features per analysis and enabled the identification of more than 1,600 phosphorylation sites using a single reversed phase separation dimension without the need for time consuming pre-fraction steps. Compared to conventional DDA LC-MS/MS experiments, a substantially higher number of peptides could be identified from a sample and this increase was more pronounced for low intensity precursor ions.

Introduction

Over the past decade, mass spectrometry (MS) has emerged as the method of choice for the identification and quantification of proteins in very complex biological samples (1). In the most widely used implementation, referred to as shotgun proteomics, protein samples are first digested, the resulting peptide mixtures are then chromatographically separated and finally sequenced by automated tandem mass spectrometry (MS/MS). Due to its conceptual and experimental simplicity, the shotgun approach has become a very popular method for the identification of proteins in a wide range of biological samples and, in combination with stable isotope labeling, also for quantitative proteomic studies (2-4). Recent technical improvements in MS instrumentation, database searching and result validation as well as advances in database annotation now make it possible to routinely identify hundreds to a few thousand of proteins in complex biological samples (5-8).

Despite this impressive progress, shotgun proteomics is not yet capable of characterizing whole proteomes and presents obvious biases, among them the discrimination against protein species of low abundance (5, 8). This is primarily a consequence of limited sequencing speed of current LC-ESI-MS/MS systems that are incapable of analyzing each precursor ion detected in complex samples together with the redundant selection of a subset of precursor ions, even if precautions like dynamic exclusion are applied. Therefore, even in repeat analyses of the same sample exhaustive identification of the low intensity precursors is not achieved (9-11).

In contrast to these approaches based on data dependent precursor ion selection (DDA), directed peptide sequencing provides the advantage of

focusing the MS/MS-analysis on non-redundant and information rich precursor ions, thereby better managing the analysis time and increasing depth of analysis (12, 13). In this regard, a two stage strategy by which all MS1 features that represent peptides are extracted from LC-MS maps and subsequently subjected to targeted sequencing, in principle, should lead to the identification of all detectable precursors (14). Since the acquisition of MS1 and MS2 spectra is naturally decoupled in MALDI-MS/MS, this platform is well suited for directed sequencing and has been applied to selectively analyze differential expression or modifications of proteins (15, 16).

The same principle is also applicable to ESI-MS, which has the potential to provide much higher sequencing speed in routine applications compared to MALDI-MS/MS. Since the peptides are not “immobilized” on the sample plate, repeat injections of the same sample are required, the first to detect the MS1 features and subsequent ones for directed sequencing. Naturally, the decoupling of feature detection and sequencing demands highly reproducible elution times and high mass accuracy. A directed sequencing strategy has been already applied on high mass accuracy ESI instruments to specifically sequence peptides of single proteins in complex mixtures (17) and for the detection of low-abundant peptide species (18). However, the number of targeted peptide ion masses was limited to a few hundred per run, a much lower number of sequencing attempts than modern ESI-MS/MS instruments are capable of performing in DDA mode. The number of targeted precursors could only be increased by sample and time-consuming multiple LC-MS/MS analyses of the same mixture.

In the present study, a high performance LTQ-FT-ICR mass spectrometer that allows segmentation of inclusion mass lists by LC elution time was used for directed sequencing. Each inclusion lists contained the mass-to-charge ratio (m/z) and elution time of the targeted precursors and was divided into segments of 3 to 5 minutes, thereby increasing the number of possible target masses to 3,000 in a one-hour LC gradient. The strategy was supported by software tools to i) automatically extract peptide features from MS1 maps and to align features over multiple LC-MS/MS patterns (13), ii) generate inclusion lists from the identified features (19), iii) control directed sequencing of the features on the inclusion list by sequential LC-MS/MS analyses of the same sample and iv) to map the MS/MS data obtained back to the initial feature list. The potential of the directed MS/MS approach was evaluated by in-depth characterization of complex peptide and phosphopeptide mixtures obtained from *D. melanogaster* lysates. The data demonstrated the high specificity and reproducibility of the method to identify a higher number of peptides from the same sample in a lower number of LC-MS/MS runs compared to standard DDA LC-MS/MS analysis, specifically in the class of low intensity precursor ions.

Material and Methods

Cell culture and phosphopeptide enrichment

All chemicals, if not otherwise mentioned, were bought at the highest available purity from Sigma-Aldrich, Taufkirchen, Germany.

Cell culture, lysis and protein digestion

D. melanogaster Kc167 cells were grown in Schneiders *Drosophila* medium (Invitrogen, Auckland, New Zealand) supplemented with 10 % fetal calf serum, 100U penicillin (Invitrogen, Auckland, New Zealand) and 100 µg/ml streptomycin (Invitrogen, Auckland, New Zealand) in an incubator at 25 °C. In order to increase the degree of phosphorylation in the *Drosophila* proteins different batches of cells were pooled which were either growing in rich medium, serum starved medium, treated for 30 min with 100 nM Rapamycin (LCIabs, Woburn, MA, USA), treated for 30 min with 100 nM insulin or treated for 30 min with 100 nM Calyculin A. Then the cells were washed with ice cold phosphate buffered saline (PBS) and re-suspended in ice cold lysis buffer containing 10 mM HEPES, pH 7.9, 1.5 mM MgCl₂, 10 mM KCl, 0.5 mM dithiothreitol (DTT) and a protease inhibitor mix (Roche, Basel, Switzerland). In order to preserve protein phosphorylation, several phosphatase inhibitors were added to a final concentration of 20 nM calyculin A, 200 nM okadaic acid, 4.8 µM cypermethrin (all bought from Merck KGaA, Darmstadt, Germany), 2 mM vanadate, 10 mM sodium pyrophosphate, 10 mM NaF and 5 mM EDTA, respectively. After 10 min incubation on ice, cells were lysed by douncing. Cell debris and nuclei were removed by centrifugation for 10 min at

4 °C using 5,500 xg. Then the cytoplasmic and membrane fraction were separated by ultracentrifugation at 100,000 xg for 60 min at 4 °C. The proteins of the cytosolic fraction (supernatant) was subjected to acetone precipitation. The protein pellets were re-solubilized in 3 mM EDTA, 20 mM TrisHCl pH 8.3 and 8 M urea. The disulfide bonds of the proteins were reduced with tris(2-carboxyethyl)phosphine (TCEP) at a final concentration of 12.5 mM at 37 °C for 1 h. The produced free thiols were alkylated with 40 mM iodoacetamide at room temperature for 1 h. The solution was diluted with 20 mM TrisHCl (pH 8.3) to a final concentration of 1.0 M urea and digested with sequencing-grade modified trypsin (Promega, Madison, Wisconsin) at 20 µg per mg of protein overnight at 37°C. Peptides were desalted on a C18 Sep-Pak cartridge (Waters, Milford, Massachusetts) and dried in a speedvac. Finally, 1 µg of peptide sample was utilized for each LC-MS/MS experiment.

Phosphopeptide isolation

Phosphopeptides were isolated using TiO₂ affinity enrichment as recently described (20). 1 µg of the phosphopeptide sample was subjected to each LC-MS/MS analysis.

Reverse phase HPLC

Peptide samples were analyzed on an Agilent 1100 microflow system (Agilent Technologies) connected to a 7 tesla Finnigan LTQ-FT-ICR instrument (Thermo Electron, Bremen, Germany) equipped with a nanoelectrospray ion source (Thermo Electron, Bremen, Germany). Peptides were separated on a RP-HPLC column (150 µm x 15 cm) packed in-house with C18 resin (Magic C18 AQ 5 µm; Michrom BioResources, Auburn, CA, USA) using a linear

gradient from 98 % solvent A (0.15 % formic acid) and 2 % solvent B (98 % acetonitrile, 2 % water, 0.15 % formic acid) to 30 % solvent B over 60 minutes (for cytosol digest) and 90 minutes (for phosphopeptide enriched samples) at a flow rate of 1.2 μ l / min.

Mass spectrometry

In DDA mode, each MS1 scan (acquired in the ICR cell) was followed by collision induced dissociation (CID, acquired in the LTQ part) of the three (for feature extraction) and five (for comparison of DDA and directed LC-MS/MS) most abundant precursor ions with dynamic exclusion for 30 seconds. Only MS1 signals exceeding 150 counts were allowed to trigger MS2 scans with wideband activation enabled. Total cycle time was approximately 1 to 1.5 s. For MS1, 10^6 ions were accumulated in the ICR cell over a maximum time of 500 ms and scanned at a resolution of 100,000 FWHM (at 400 m/z). MS2 and MS3 spectra were acquired using the normal scan mode, a target setting of 10^4 ions and accumulation time of 250 ms. Singly charged ions and ions with unassigned charge state were excluded from triggering MS2 events. The normalized collision energy was set to 30%, and one microscan was acquired for each spectrum. For phosphopeptide analysis, the mass spectrometer automatically switched between MS, MS2, and neutral loss-dependent MS3 acquisition. Data-dependent settings were chosen to trigger an MS3 scan when a neutral loss of 97.97, 48.99, 32.66, 24.5 or 19.6 Da was detected among the ten most intense fragment ions.

Peak detection

First, the data of the initial three LC-MS (mapping) runs (raw format) was converted to the profile mzXML format (21). Then, the in-house developed software system *SuperHirn* (22) was used for i) detection, ii) de-isotoping, iii) peak integration and iv) alignment of detected features over multiple LC/MS patterns. Peak intensities were measured by calculating peak areas from extracted ion chromatograms (XICs) of each MS signal. Highly stringent criteria were applied to filter the detected peaks for peptide signals. Specifically, the algorithm searches for peaks patterns matching isotope distributions typical for peptides within the m/z value range under investigation. Peaks had to be detected in at least two subsequent MS1 scans and with a minimum of three isotopic peaks to be considered. Only peaks that could be found in at least two LC-MS runs were considered and singly charged masses were excluded. Finally, a list of the relevant features was generated and used to build mass inclusion lists for directed MS-sequencing.

Generation of inclusion lists

To make the generation of inclusion lists less time consuming, less prone to human error and easier to reproduce we developed the “Inclusion List Builder” software. The Inclusion List Builder has a rich graphical user interface and is implemented as a plug-in for the *Prequips* platform (19) (Download: http://tools.proteomecenter.org/wiki/index.php?title=Software:Prequips:Inclusion_List_Builder).

The table containing all features extracted from initial LC-MS runs is imported through the Prequips’ data provider interface and converted into a so-called

“Master Table” by the Inclusion List Builder. The master table contains the m/z ratios, retention times, averaged peak areas and charge states for all features identified by the *SuperHirn* algorithm. Through interactive application of filters to feature attributes inclusion lists are created as subsets of the master table and segmented by retention time. This is necessary, since the number of features present in the table usually exceeds the number of possible sequencing cycles that the mass spectrometer can acquire in a single run. After segmentation the inclusion lists are exported as tables (.csv file format) that can be read by the MS-instrument software.

In the study presented here, the following software settings were used for the directed analysis of the peptide mixtures. The 9,680 features extracted from the three map runs analyzing the drosophila lysate digest were split by their intensity into five bins, each consisting of 2,000 masses (1,680 for the last bin containing the least intense features) using the following average peak area thresholds; very high: $2.6 \times 10^9 - 3 \times 10^7$, high: $3.0 - 1.4 \times 10^7$, medium: $1.4 \times 10^7 - 8.1 \times 10^6$, low: $8.1 - 4.9 \times 10^6$, very low: $4.9 \times 10^6 - 5.3 \times 10^5$. Each subset was further clustered into 3 or 5-minute segments using the following elution time values (in minutes); 0 - 27.5 - 32.5 - 37.5 - 40.5 - 43.5 - 46.5 - 49.5 - 52.5 - 55.5 - 58.5 - 61.5 - 66.5 - 71.5 - 80. The start and stop time of each 5 (3) minute time segment was extended by 2.4 (1.4) and 2.5 (1.5) minutes, respectively, to compensate for variations in retention time. Since the instrument software requires a few seconds to load/delete the new/old masses in each time segment, a delay of six seconds was implemented for each start time. For directed analysis of the phosphopeptide-enriched sample (90 minute gradient), the features were clustered using the following elution

time bins: 0 - 22.5 - 27.5 - 32.5 - 37.5 - 42.5 - 47.5 - 52.5 - 57.5 - 62.5 - 67.5 - 72.5 - 77.5 - 82.5 - 87.5 - 95 - 110. In summary, this setup allowed, assuming equal distribution, the inclusion of up to 3,000 features in a single directed LC-MS/MS analysis using a 60 minute gradient. It is important to note that applying shorter time segments could further increase this number.

Directed MS-sequencing of features

The generated inclusion lists (.csv file format) were directly imported into the global mass list parent ion table of the MS operating software (XCalibur 2.0 SR 1, Thermo Electron, Bremen, Germany) and activated. Basically, the settings for targeted LC-MS/MS were similar to those described above with a few modifications. First, the dynamic exclusion mass window that is also setting the m/z tolerance for the inclusion list masses was narrowed to ± 10 ppm for all directed analyses with enabled monoisotopic precursor selection and to ± 5 ppm when this option was turned off. Ion signals for which no charge could be assigned were also allowed to trigger MS2 scans. The dynamic exclusion time was reduced to 10 seconds to acquire multiple MS2 scans for each feature. For directed sequencing in preview off mode, this option was disabled and the resolution of the MS1 scan in the ICR cell reduced to 50,000. For sequencing low abundant features of low abundance, the monoisotopic precursor selection option was disabled and, to minimize unspecific sequencing, the threshold required for triggering MS2 events was raised from 150 to 3,000 counts.

Database searching

All acquired MS2 and MS3 spectra were searched against the *Drosophila* Flybase protein database (*D. melanogaster*, release 4.3; Mar 2006; 19645 entries) that also contained the protein sequence of bovine trypsin and human keratins, using the Bioworks (Version 3.2) software (Thermo Electron, San Jose, CA). The search criteria were as follows: full tryptic specificity was required (cleavage after lysine or arginine residues, unless followed by proline); 2 missed cleavages were allowed; carbamidomethylation (C) was set as fixed modification; oxidation (M) and, if required, phosphorylation (STY) were applied as variable modifications; mass tolerance of the precursor ion and the fragment ions was 10 ppm and 0.8 Da, respectively. In addition to this, each data set was searched against a decoy Flybase protein database (Version 4.3), as described previously to assess the number of false positive peptide identifications (23). Based on this approach, the error rate was set to a maximum of 1% (error rate = 2 x percentage of decoy hits) using the following thresholds: Peptide probability (prob) < 0.01; Final score (Sf) > 0.2 for unphosphorylated and > 0.4 for phosphorylated peptides. For each peptide sequence identified, all matching gene numbers (Flybase gene ID (FBgn)) and protein accession entries (Computed gene (CG)) were determined and displayed. In addition to this, Occam's Razor logic as implemented in Protein Prophet (24) was applied to calculate the number of identified proteins (CG entries). In brief, redundant protein entries were removed by clustering peptides matching to multiple members of a protein family to a single protein group and considered as a single identification. Furthermore, when multiple

proteins shared a peptide sequence, it was only assigned to the protein identified with the highest number of peptide assignments. In-house software was used for the calculation of non-redundant phosphorylation sites present in the data sets obtained as recently reported (25).

For comprehensive intensity comparison of the peptides identified by DDA and directed LC-MS/MS analysis (Figure 3B), the MS1 signal intensities were determined using the Bioworks software, to also include peptides exclusively identified in DDA mode. Therefore, the peak height of every single peptide precursor ion was determined by applying the following parameter: mass range of 0.03 Da; intensity threshold of minimal 1,000 and 3 smoothing points allowed. The intensity of the same peptide ions detected in multiple runs was averaged.

Assignment of identified peptides to inclusion lists

An Excel table (in tab-delimited format) containing the search results from the directed LC-MS/MS runs was imported into the *Prequips* software. Spectral data in mzXML format was loaded to obtain access to the corresponding retention times. All identified peptides were mapped to one or more features in the master table through the MS2 scans associated with them.

For the mapping algorithm we define a scan as the triple (corrected retention time t'_R , feature at mass m and charge z), where the feature mass m is defined as

$$m = (\text{precursor neutral mass} + (z * 1.00727)) / z$$

and the corrected retention time t'_R is computed as

$$t'_R = a * t_R + b$$

from the measured retention time t_R . Since all LC-MS/MS experiments were performed using the same C18 column resulting in minimal shifts in feature elution times, no corrections of t_R were required. Therefore, a was set to 1 and the value for b was set 0. The following algorithm was applied for each identified peptide p :

1) For every scan s associated with p find all features f in the master table that fulfill each of the following conditions:

$$t'_R(s) - \tau \leq t'_R(f) \leq t'_R(s) + \tau$$

$$m(s) - \mu \leq m(f) \leq m(s) + \mu$$

$$z(s) = z(f)$$

The retention time tolerance τ (in minutes) and the mass tolerance μ (in ppm) define windows of 2τ and 2μ centered on $t'_R(s)$ and $m(s)$, respectively.

2) Map p to all f fulfilling the conditions.

A single peptide can be mapped to the same feature multiple times if it is associated with more than one scan. Also, more than one distinct peptide can be mapped to a feature. The peptide with the lowest retention time and mass difference is automatically selected as the best hit. The investigator can manually assign any other of the peptides mapped as the best hit for a given feature if there is supporting evidence.

For mapping back identified features to the initial inclusion list, the theoretical m/z values and the elution time of the identified peptides was matched with the precursor masses and elution times in the inclusion list. A mass tolerance of 0.01 Da and a time tolerance of 1 minute were allowed. Since no pre-

column was used during LC, very hydrophilic peptides, which were not or only partially retained by the C18 resin, showed very inconsistent retention times. Therefore, all peptides having LC retention times of less than 30 minutes were matched only by their m/z and charge values.

Results

The directed precursor ion selection workflow

The general workflow of the directed LC-MS/MS analysis described in this manuscript is outlined in Figure 1. It consists of the following steps:

(1. Sample preparation) The proteins in a sample obtained from any source (cells, tissue, body fluids, etc.) are reduced, alkylated and enzymatically cleaved with trypsin.

(2. Feature detection) Sample aliquots are then analyzed by high performance LC-MS/MS whereby the MS1 features are recorded and selected precursor ions are picked for CID by DDA. Usually, we performed three independent LC-MS/MS runs, each consuming 1 μ g of total peptide mass. The data obtained is subsequently analyzed offline by the in-house software tool *SuperHirn* (13) that automatically extracts all detectable MS1 signals from the individual LC-MS patterns by applying peak detection, noise reduction, de-isotoping and intensity determination algorithms to each detected peak. The detected ions are then aligned within the respective patterns.

(3. Master table generation) The software overlays over all detected peaks an isotope distribution template that is typical for peptides, thereby discriminating between peptide and contaminant signals, and finally generates a master

table of all masses that are very likely to be peptide ions. Each of these features is described by its m/z ratio, charge state, intensity and chromatographic retention time. All features are then imported into a second in-house software tool, *Prequips* that supports data filtering, generation of inclusion list files and mapping MS2 data back to the respective MS1 features in an automated manner (19). The alignment of features over multiple patterns was a very effective method to reduce noise or signals from non-peptidic material if only MS1 signals that are found in at least two of the three replicate LC-MS runs were considered for further analysis. In the end, around 10,000 features are detected in a typical experiment.

(4. Inclusion list creation) Since for most samples the number of features detected exceeded the number of available sequencing cycles, the feature list was further divided according to the sequencing speed of the MS instrument used. For the LTQ-FT-ICR instrument employed in our study, a list of 10,000 features was divided into five LC-MS/MS inclusion lists of 2,000 features each. In general, the features were grouped into segments of five (three for highly populated areas) minutes and flanked by 2.5 (1.5) minutes extra time, respectively, resulting in segments of 10 (6) minutes, respectively. Hence, each feature could tolerate a shift in retention time during LC of at least 2.5 (1.5) minutes. Initial tests showed that the current instrument control software (Xcalibur 2.0 SR1) limited the number of ions on an inclusion list to a maximum of 500. Thus the construction of the inclusion list required that the number of features from two overlapping segments did not exceed 500. Besides, the lower number of features early and late in the gradient allowed the generation of longer segments with a higher time tolerance in those

regions. The long time segment at the beginning is essential for targeting hydrophilic peptides, because these molecules do not, or only weakly, bind to the RP-LC column and consequently showed considerable shifts in elution times between different runs. For samples that showed a common feature distribution, a maximum of 3,000 features could be sequenced per hour (~ 1 feature / sec). It is worth mentioning that this number could be further increased by a further reduction of the time segments.

(5. Feature identification) The MS2 data obtained from the directed LC-MS/MS analyses was searched against protein databases and, together with the corresponding mzXML files, mapped back to the master table using the *Prequips* software.

Reproducibility of LC-MS

The success of an inclusion list experiment strongly depended on the reproducibility of both, LC and MS performances. Figure S1 shows that variations in retention time were below 20 seconds and mass accuracy was better than 2 ppm. These variations are well within the time and m/z tolerances of 2.5 minutes and 10 ppm, respectively, used for directed MS2. Naturally, the time and m/z tolerances should be kept as small as possible to minimize random MS-sequencing events.

Optimization of MS-parameters for improved identification of features

To optimize the number of features identified in a targeted precursor ion selection experiment we generated a peptide mixture from *D. melanogaster*

Kc167 cells and subjected aliquots to repeat analyses in which relevant experimental parameters were varied.

Cells (10^8) were harvested, disrupted and the cytosolic fraction was isolated. The proteins were alkylated, trypsinized and three aliquots, each containing 1 μ g of peptides, were analyzed as described above (Figure 1). A total of 9,680 unique features could be detected (see Table S3) that were imported into the master table. The features were grouped by decreasing intensity into five inclusion lists each containing around 2,000 features, and subjected to directed LC-MS/MS analysis.

Subsequently, the influence of two important parameters for triggering MS2 in the LTQ-FT MS employed, the preview mode and the monoisotopic precursor selection (MPS), respectively, was evaluated. With preview mode on (default mode), the instrument acquires a low resolution MS1 spectrum (pre-scan) that takes around 20% of the total scan time and continues scanning the ions for high resolution while the LTQ is performing MS2 scans of selected ions in parallel (Figure 2C). This significantly increases MS2 scan rates but peptide ions for MS2 are selected from low resolution MS1 scans. By disabling the preview mode and reducing the resolution to 50,000 for MS1, the mass accuracy of the peptide ions selected online for directed MS2 could be improved. Consequently, 8% (13%) more features were identified (sequenced) from the Kc167 cell sample from the five inclusion lists with preview mode disabled (Figure 2A, Table S1, S4, S5). Despite the slightly longer cycle times required (Figure 2D), the speed of the MS instrument used was still sufficient to selectively sequence up to 2,000 features in a 60 minute gradient (Table S1). It is important to note that the improvements were

particularly pronounced for features with lower intensities. Additionally, the MS2 selection efficiency of peptide ions of low abundance could be further improved by disabling MPS, which makes every single isotopic peak in the MS1 scans accessible for triggering MS2. As a consequence, the yield of identified low abundant peptide signals could be further improved from 33% to 40% and 26% to 35% for low and very low intense features, respectively (Figure 2A).

It is worth mentioning that increasing the number of ions into the ICR-cell for MS1 scans also slightly increased the numbers of identified low abundance ions, however, to a lower extend than by the disabling MPS (data not shown). This is due to the fact that high ion numbers in the ICR-cell lead to space charge effects that lower peptide mass accuracy (26, 27). Therefore, some features might fall out of the m/z tolerances applied and would not be sequenced if the ICR cell is filled with a high number of ions.

Using the optimized MS-parameters, around 80% of all detected features were selected for sequencing and 48% of those could be confidently identified, resulting in the identification of 3,931 unique peptides and 793 non-redundant gene products (Table S1+S6). In addition, the occurrence of random sequencing of MS-signals not selected for directed MS2 was very low since only about 11.6% (516) of all identified peptides in the five inclusion list runs could not be mapped back to the master feature list. This demonstrates that the whole process starting with feature extraction from high resolution MS1 maps followed by directed sequencing using an inclusion list protocol and mapping back the MS2 data to the features is very effective and specific.

Comparison of data dependent and directed LC-MS/MS

To assess the performance of the directed LC-MS/MS approach in comparison to CID experiments using DDA, we compared the data obtained from applying the protocols optimized as described above with the results of five repeat LC-MS/MS runs using DDA. The sample was a tryptic digest of Kc167 cell cytosolic fraction and in each analysis 1 μ g of total peptide mass was injected. To increase the number of MS2 spectra acquired, the instrument was programmed to randomly select for CID the five most intense precursor ion signals detected in a survey scan in the LC-MS/MS experiment with DDA and the peptides identified by either strategy were compared.

As shown in Figure 3A (blue line), 2,493 unique peptides were identified in the first DDA LC-MS/MS run, but then the contribution of additional LC-MS/MS runs of the same sample to the overall number of peptides was decreasing rapidly, eventually identifying only 1,083 (43.4%) additional peptides in the four subsequent LC-MS/MS runs (Table S2+S7). This effect was even more apparent at the protein level where only 111 (21.0%) new gene products could be discovered by the additional four LC-MS/MS experiments. Notably, the number of identified peptides over all five DDA runs was highly consistent, ranging from 2,461 - 2,493 (Table S2+S7). This clearly demonstrates a strong sequencing redundancy in shotgun LC-MS/MS, which is in agreement with observations obtained from other recent studies (9-11).

In contrast, the directed approach presented here allowed the non-redundant sequencing of every single feature in only one LC-MS/MS run and therefore, the degree of novel peptide identifications in each analysis was much higher,

resulting in a much steeper curve (Figure 3A, orange line). Here, using the optimized MS-parameters shown above (Preview mode and MPS disabled, Figure 2), 2,746 (237%) additional peptides could be determined after the first directed LC-MS/MS run resulting in a total of 3,931 unique peptide identifications. Using non-redundant sequencing also allowed the identification of a significant number of additional proteins in each directed run compared to DDA LC-MS/MS (Table S1). Eventually, a higher number of peptides and proteins were identified in five directed runs than in five DDA LC-MS/MS analyses, despite the much lower number of sequencing events, clearly indicating the higher information content of the ions extracted offline over DDA LC-MS/MS (Table S1 + S2). Interestingly, the INL curve (Figure 3A) is only slightly flattening with decreased feature signal intensity, which suggests that potentially even more peptides can be identified by the directed approach if less stringent peak picking criteria would be applied and additional inclusion list runs would be carried out on the same sample.

It is important to point out that current mass spectrometers offer possibilities to sequence additional low-abundant peptides in DDA mode (28). However, most of them drastically reduce the number of MS1 scans acquired, and therefore lower reliable quantification of complex peptide mixtures. Conversely, directed LC-MS/MS enables the identification of low-abundant peptides without affecting MS1 performance and thus quantification accuracy.

Combination of DDA and directed LC-MS/MS

As shown in Figure 3A, three inclusion lists were necessary to identify as many peptides/proteins as in a single DDA LC-MS/MS experiment. Therefore,

excluding already selected features from the initial LC-MS run for the generation of MS1 maps and specifically targeting the remaining features would be expected to reduce the number of directed LC-MS/MS runs required to sequence every detectable feature. We applied a combined strategy of two initial DDA runs with subsequent directed runs to the Kc167 cell sample described above. As shown in figure 3A, the results of more than 5,000 sequencing attempts of the first two DDA LC-MS/MS analyses runs could be mapped back to the feature list, thereby reducing the number of features from 9,680 to 4,323 that were specifically targeted in two additional LC-MS/MS experiments. The first directed run already identified as many novel peptides as the three additional DDA runs together and eventually, 4,273 unique peptides corresponding to 804 different proteins could be identified after only four LC-MS/MS runs (Figure 3A, green line, Table S8). In total, a higher number of peptides could be identified with less analytical effort and time by the combined approach than by the DDA or directed approach alone. Notably, the high performance hybrid LTQ-FT mass spectrometer used for our experiments is very well suited for the combined approach, since the ICR cell enables the acquisition of high resolution MS1 data while, in parallel, the LTQ part can collect MS2 data without sacrificing time, resolution or sensitivity for MS1 data acquisition (Figure 2C).

To evaluate the intensity distribution of the peptides identified by both, DDA alone (Figure 3A, blue line) and in combination with directed (Figure 3A, green line) LC-MS/MS, the precursor ion abundances of these peptides were calculated and compared. To obtain an un-biased intensity determination of

all identified precursor ions, their base peak intensities were calculated using the Bioworks software tool. As shown in Figure 3B, most of the peptide ions identified in DDA mode had intensities between 10^6 and 2×10^6 counts (Figure 3B, blue line). By contrast, the peak maximum of the distribution of peptides exclusively identified by the two inclusion list runs is clearly shifted towards precursor ions with lower abundance. The apex of the distribution of the peptides determined by the inclusion list in the combined approach was around 2×10^5 to 4×10^5 and thus about 5-times less intense than that of the DDA alone strategy (Figure 3B, red line). In combination with the data obtained from the first two DDA runs, the intensity of peptides determined by the combined approach was much less biased towards high abundant signals and more equally distributed (Figure 3B, green line). This clearly indicates that in complex peptide mixtures, a higher number of peptide ions with low intensities could be identified by directed than by DDA LC-MS/MS approaches.

Reproducibility of directed LC-MS/MS sequencing

To evaluate the reproducibility of the directed approach for both, feature selection and identification, 200 identified features from each of the five intensity groups (very high to very low) were combined and re-analyzed five times. Overall, the selected features spanned more than three orders of magnitude in signal intensity. The data are shown in Table 1 and Figure S2 . Of the 1,000 features targeted, more than 85% did trigger sequence attempts and around 70% could be confidently identified in all five LC-MS/MS runs. Furthermore, of the 1,000 features on the respective inclusion list, 984 did

trigger a MS2 scan event and 945 could be assigned to the correct peptide sequence in at least one run. Since the directed approach requires each feature to be detected online, the selection of features for sequencing strongly depends on their signal intensity. Whereas all of the 200 features in the group with the highest signal intensities were selected for MS2 in all five repeat LC-experiments, that number decreased to 148 (74%) in the group with the lowest signal intensities (Figure S2B). A similar trend was observable for the number of correctly identified peptides albeit with a steeper decline at lower feature intensities. However, the MS2 spectra obtained can be useful for side-by-side comparison of correctly assigned spectra to confirm the feature identity (29), even if the spectral quality is not sufficient for the assignment of a peptide sequence by a database search engine.

Directed analysis of a complex phosphopeptide mixture

Phosphopeptides, specifically those phosphorylated at serine or threonine residues are notoriously difficult to identify, due to their specific fragmentation patterns in ion trap instruments.

To evaluate the performance of directed sequencing for the identification of phosphopeptides we applied the combined directed approach described above (Figure 3A, green line) to a sample consisting of phosphopeptides isolated by TiO₂ affinity chromatography (20, 30) from a cytosolic fraction of Kc167-cells. The sample was separated using a RP-LC gradient that was extended by 30 minutes compared to the one used for non-phosphorylated peptides to compensate for the longer MS acquisition times applied for phosphopeptide analysis (MS2 followed by MS3 of the neutral loss peak of

-98 Da corresponding to the loss of phosphate). To maximize the number of identified features for a specific number of LC-MS runs, the experiment was designed such that in every run a different set of peptides was selected. Specifically, the following protocol was applied: First, the sample was analyzed by DDA LC-MS/MS, selecting precursor ions at $[M+2H]^{2+}$ in the first run and ions at charge states higher than $[M+2H]^{2+}$ in the second run, respectively. Then, all MS1 peaks were extracted from the high resolution MS1 maps obtained in those first two runs and filtered as described above. A total of more than 7,776 features that aligned over both runs were detected (Table S9), about half of which were already sequenced by the two initial LC-MS/MS runs and excluded from following directed LC-MS/MS. After generating a series of inclusion lists, the remaining 4,000 features were subjected to directed sequencing in two additional LC-MS/MS experiments. To further increase the number of detected phosphopeptides, the 2,000 most abundant features showing a neutral loss peak but could not be confidently assigned to a peptide sequence were re-analyzed in one additional LC-MS/MS run using a different MS-sequencing method especially designed for phosphopeptide analysis (31).

Every directed LC-MS/MS run identified a considerable number of previously unidentified peptides. In total, after five LC-MS/MS runs, more than 1,600 phosphorylation sites could be identified (Table 2 + S10, Figure S3). Interestingly, 1,500 (87.2%) of the 1,721 identified peptides carried at least one phosphate group confirming a high specificity for phosphopeptides of the TiO_2 affinity enrichment. Of the 1,628 phosphorylation events detected, the exact site of phosphorylation of 1,204 sites could be determined with a

probability of more than 90% ($\Delta\text{CN} > 0.1$) (32). The distribution of phosphorylated amino acids was similar to that in other studies (20, 33) with most sites found on serine (82.7%), followed by threonine (15.4%) and tyrosine (1.9%) residues. In addition to this, the majority of phosphopeptides identified contained one phospho group (92.1%) whereas only 110 (7.3%) and 9 (0.6%) were phosphorylated on two or three different residues, respectively. More importantly, compared to the DDA runs, 65% additional protein phosphorylation sites were identified by the three directed LC-MS/MS runs (run 3-5) of which 107 were identified by re-analyzing features that showed a neutral-loss of phosphate (-98 Da) during CID (run 5). Compared to a single DDA LC-MS/MS analysis of this sample that detected a total of 720 phosphorylation sites (Table S11), the directed strategy employed here enabled the identification of a 2.3-times higher number of phosphosites after five LC-MS/MS runs. Annotated spectra of all phosphopeptides identified by the directed approach and by the single DDA LC-MS/MS run are shown in the Supplementary Figures 4 and 5, respectively.

The value of the additional information obtained with directed LC-MS/MS can be demonstrated by the increased protein phosphorylation coverage of the Wnt signaling pathway, which is implicated in the genesis of cancer (34, 35). Table 3 shows all 8 identified phosphoproteins and their 13 phosphorylation sites that could be assigned to this pathway using PANTHER (36). Whereas 5 phosphoproteins and 7 phosphorylation sites could be detected by the two initial DDA LC-MS/MS analyses (1-2), the three following directed LC-MS/MS runs (3-5) exclusively detected 3 phosphoproteins and 6 phosphorylation sites that increased the overall coverage of the pathway by 60% and 85.7%,

respectively. For example, phosphorylation of ATP-dependent Helicase SWR1 (Gene name: CG5899) at serine 169, 172 and 841 could be determined with high confidence only by additional directed LC-MS/MS. Even though the precursor ion signal intensity for fragment ion spectra acquired by directed sequencing were generally lower than those of the DDA runs, their quality was high. For instance, the complete y-ion series, with the exception of y1 and y2, of the doubly phosphorylated peptide “K.DQVYDpSDDpSDSEMSTK.M” could be assigned from its fragment spectrum (Figure 4B). Interestingly, the precursor ion was not picked for fragmentation in the first two runs although it is only 5-times less abundant than the highest peak present in the corresponding MS1 spectrum (Figure 4A). Obviously, the sequencing speed of the MS was not sufficient to acquire MS2 spectra from the high number of co-eluting peptide ion present in the MS1 spectrum. This agrees very well with our results obtained from the sample consisting of non-phosphorylated peptides (Figure 3B).

Discussion

This study describes a reproducible, sensitive and integrated computational and mass spectrometric method for directed sequencing of a high number of peptide ions within complex mixtures. In contrast to concomitant MS1 and MS2 data acquisition during DDA ESI LC-MS/MS, the directed approach described decouples MS1 and MS2 spectra collection. In a first step, potential peptide ion signals are extracted offline from the pattern generated in initial LC-MS/MS runs. For this task we developed the software tools *SuperHirn* and

Prequips. In a second step, such features were subjected to directed sequencing in subsequent LC-MS/MS runs and the identified features automatically mapped back to the list of previously detected features. As we make the software tools developed for this method publicly available and as it uses generic data formats (21, 24, 37) for which converters for various MS-instruments are available (<http://tools.proteomecenter.org/wiki/index.php?title=Formats:mzXML>), the presented approach is applicable to any high performance MS platform that allows direct sequencing by inclusion list. It is important to point out that high resolution scans are required only for MS1 data acquisition and not for MS2 sequencing. Notably, hybrid MS instruments, like the LTQ-FT used in this study are preferred, since they provide the unique advantage of parallel MS1 and MS2 spectra acquisition. Therefore, MS2 data can already be obtained in the initial LC-MS/MS runs with minimal impairment of the speed and sensitivity for acquiring high quality MS1 maps. Compared to current DDA LC-MS/MS, the described directed strategy provides several advantages.

First, the offline feature extraction facilitates the identification of peptides with low intensity precursor ion signals. Whereas peak selection in DDA LC-MS/MS analysis is based on one single MS1 scan, offline peptide ion detection allows summing up all isotopic signals of an eluting peptide ion. This simplifies the selection of potential peptide ions based on the distribution of isotopic clusters as well as the determination of charge state and monoisotopic precursor mass. Moreover, the alignment of detected peptide ions over multiple runs helps to distinguish between peptide derived signals

and chemical and electronic noise. This significantly improved the generation of a feature list with a high content of peptide ions. As shown above, the directed sequencing of these information rich features resulted in a higher number of unique peptides identified compared to online CID by DDA. It is worth mentioning that the number of MS2 scans acquired by the directed approach was 3-times lower than by DDA LC-MS/MS, which considerably reduced the efforts for subsequent data analysis.

Second in directed LC-MS/MS, the CID parameters can be adjusted and optimized according to the feature properties. For instance, a higher number of very intense features could be identified in one LC-MS/MS experiment by applying shorter ion accumulation and scan times whereas longer gating and scan times were employed for low abundant peptide ions. It is important to point out that the directed approach offers the opportunity of re-analyzing all features that could not be assigned to a peptide sequence in the first analysis using optimized MS-parameters. As shown by the results obtained in this study, more than 100 additional phosphorylation sites were determined by iterative analysis of phosphopeptides that showed a neutral loss peak but could not be confidently identified on the basis of their MS2 or MS3 spectra using the multistage activation mode for MS/MS data acquisition (31). The application of optimized CID parameters for each feature not only allows for the identification of more peptides but can also be employed to confirm peptide ions with questionable identity.

Third, the directed approach offers the possibility of selecting for CID precursor ions with specific properties. These include the charge state of a peptide that is most likely to yield an informative fragment ion spectrum, particular patterns after stable isotope labeling (13, 15, 16, 38), characteristic isotope distributions generated by tagging selected functional groups with suitable reagents (39), both isotopic version of cross-linked peptides (40) or redundant identification of peptides of interest over multiple samples (e.g. time course experiments) (13).

For analyzing complex mixtures, the most important improvement of directed versus DDA LC-MS/MS results from the fact that the directed approach copes with the problem of “undersampling” during LC-MS/MS analysis meaning that not all precursor ions present in the MS1 scans can be sequenced using DDA based LC-MS/MS (8-11). This results in the preferred identification of peptides of high abundance (Figure 3A). Conversely, using the directed approach, the sequencing speed of the MS instrument used is no longer a limiting factor with respect to the under sampling problem virtually allowing any detectable peptide ion in a sample to be MS-sequenced. Indeed, a higher number of peptides were identified by using the inclusion list based approach with most having a precursor intensity five times lower than the majority of peptides determined by the DDA strategy. The exclusive identification of low intensity peptide ions by directed LC-MS/MS was particularly pronounced for MS1 scans containing large numbers of peptide ions, clearly indicating incomplete sequencing by online DDA LC-MS/MS due to the limited MS-sequencing

speed and a more thorough sequencing of precursor ions by the directed approach.

Nonetheless, the number of peptides identified by directed LC-MS/MS decreased rapidly with lower MS1 signal intensities. This can be ascribed to the fact that very low intense peaks do not reach the detection limit in all replicate runs, show higher mass tolerances and are consequently difficult to detect and to align across multiple maps. A higher dynamic range in the MS1 scans would be desirable, however, this is limited by the loading capacity of the ICR cell (26, 27). Certainly, the dynamic intensity range of detected peptide ions can be extended by additional time-consuming sample fractionation steps (5, 41), which can be employed in combination with the directed approach. Second, and more importantly, each feature needs to be detected online by the MS-instrument to trigger an MS2. As shown above, changing the MS-detection parameters increased this number, but for a considerable fraction of low abundance features detected offline, no MS2 scan was acquired. Therefore, further improvements in online monoisotopic peak detection of peptide derived precursor ions could definitely increase the number of CIDs for low abundant peptide ions.

In conclusion, directed sequencing of non-redundant, high quality features enables the identification of a higher number of peptides with less analytical effort than current DDA LC-MS/MS based methods. For instance, more than 1,600 phosphorylation sites could be identified using a single dimension reversed phase separation without the need for time-consuming sample pre-fraction steps. The implemented software tools used are freely available and

compatible with generic, public accessible data formats and therefore applicable to most high performance LC-MS (MS1-level) platforms. With the increasing availability of reproducible high performance LC-MS/MS systems and its capability to identify features of interest specifically and redundantly over a wide intensity range, the directed approach presented here is well suited for in-depth and high throughput characterization of complex protein samples and will find wide application in future LC-MS/MS based proteome studies.

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Figure legends

Figure 1: **Workflow of directed LC-MS/MS analysis.** Aliquots of a complex peptide mixture are analyzed by LC-MS/MS and all MS1 peaks are extracted. Subsequently, sub-sets of features are selected and specifically sequenced by directed LC-MS/MS until MS/MS data is obtained for all detected features. (see text for details).

Figure 2: **Evaluation of different MS-conditions for improved feature sequencing.** The 9,680 features extracted from the cytosolic digest from the *D. melanogaster* Kc167 cell line were split into five bins based on their intensity, each containing around 2,000 masses and subjected to directed LC-MS/MS using a LTQ-FT MS. The percentage of (A) sequenced and (B) identified features in each experiment using different MS-parameters is shown. (C) Parallel acquisition of a high resolution MS1 scan in the ICR cell at a resolution of 100,000 and three MS2 spectra in the LTQ in Preview ON mode. (D) Sequential MS1 and MS2 data acquisition using a lower resolution in the ICR cell (50,000) in Preview OFF mode. The time required for each MS cycle is indicated. Additional experiments for features with low and very low intensity were performed using Preview OFF mode and disabling monoisotopic precursor selection (Preview + MPS OFF).

Figure 3: **Comparison of data dependent and directed LC-MS/MS analysis.** (A) Unique peptides identifications of the cytosolic digest from the *D. melanogaster* Kc167 cell line obtained with multiple data dependent LC-MS/MS experiments (DDA, blue line), optimized directed approach described

in Figure 2 (INL, orange line) and a combination of both methods (Combined, green line). Here, two DDA LC-MS/MS experiments were carried out and only extracted features without MS2 data subjected to two additional directed LC-MS/MS analyses. (B) Base peak intensity distribution of the peptide precursor ions identified using either DDA mode (DDA, blue line) or the combined approach shown in A (Combined, green line). The abundance profile of the peptides exclusively detected with the two directed LC-MS/MS runs in the combined approach is also indicated (INL (from combined), red line).

Figure 4: **Determination of two phosphorylation sites in ATP-dependent Helicase SWR1 (Gene name: CG5899) by directed LC-MS/MS.** (A) MS-spectrum showing the feature selected for sequencing during directed LC-MS/MS data acquisition. (B) Fragmentation pattern (MS/MS-spectrum) obtained by collision-induced dissociation of the phosphopeptide precursor indicated in A. The sequence of the identified phosphopeptide, including the two phosphorylated serine residues (pS), is shown. Fragmentation of the peptide backbone during tandem mass analysis would result in the characteristic b- and y-ions displayed. All detected fragment ions (bold) including loss of phosphoric acid (*) are indicated.

Supplementary Figure Legends

Figure S1: **Mass and elution time reproducibility during LC-MS analysis.**

(A-C) Base peak chromatograms obtained from three repetitive LC-MS analyses of aliquots of a cytosolic digest from the *D. melanogaster* Kc167 cell line. The retention times of the highest peaks are indicated. (D-F) Representative MS1 scan acquired in the ICR cell at a resolution of 100,000 FWHM (at 400 m/z) showing several features detected at a retention time of 50.41 min in each LC-MS run (A-C).

Figure S2: **Reproducibility and robustness of directed feature sequencing.**

An inclusion list comprising 200 identified features from each of the five intensity groups shown in Figure 2B was generated and re-analyzed five times by directed LC-MS/MS. (A) Number of times a features was sequenced or identified in the five replicates. (B) Number of features sequenced and identified in all five replicate runs based on their intensity.

Figure S3: **Non-redundant, directed LC-MS/MS analysis of a**

phosphopeptides mixture. A cytosolic digest from the *D. melanogaster* Kc167 cell line enriched for phosphopeptides using TiO₂ affinity chromatography was analyzed by the combined directed approach shown in Figure 3A. The first two LC-MS/MS runs (1 (DDA 2+) and 2 (DDA >2+)) were performed in DDA mode, allowing only doubly or triply and higher charged peptide ions, respectively, to trigger MS2 scans. Extracted features obtained from these to runs without MS2 data were subjected to two additional directed LC-MS/MS runs (3 (INL-1) and 4 (INL-2)). In run 5 (INL-NL), only precursor

ions that showed a neutral loss peak in the acquired MS2 spectra but could not be confidently assigned to a peptide sequence were re-sequenced with enabled multi stage activation mode in the LTQ-FT MS. The total number of identified peptides, phosphopeptides and phosphorylation sites together with the number of LC-MS/MS runs carried out is indicated.

Table 1 : Reproducibility of directed LC-MS/MS

# of observations	Feature Intensity*	# of features targeted**	# of features sequenced	# of features identified
1	All	1000	984	945
2	All	1000	972	906
3	All	1000	953	863
4	All	1000	923	803
5	All	1000	865	695
5	Very high	200	200	192
5	High	200	186	164
5	Medium	200	170	129
5	Low	200	161	117
5	Very low	200	148	93

*) Feature intensity bins based on feature area determined by SuperHirn (very high: 2.6×10^9 – 3×10^7 , high: 3.0 – 1.4×10^7 , medium: 1.4×10^7 - 8.1×10^6 , low: 8.1 – 4.9×10^6 , very low: 4.9×10^6 – 5.3×10^5)

**) The same 1000 features were targeted in each run, including 200 features of each intensity bin

Table 2: Identified phosphopeptides using directed LC-MS/MS

LC-MS/MS run*	# of peptides identified**	# of phosphopeptides identified**	# of phosphorylation sites identified**	% of phosphopeptides identified**
1 (DDA 2+)	742	655	678	88.27
2 (DDA >2+)	1085	949	997	87.47
3 (INL-1)	1439	1254	1347	87.14
4 (INL-2)	1621	1405	1521	86.67
5 (INL-NL)	1721	1500	1628	87.16

*) Run 1 (DDA 2+) and 2 (DDA >2+): only doubly or higher charges peptide ions, respectively, were allowed to trigger CID by DDA. Run 3 (INL-1) and 4 (INL-2): directed LC-MS/MS of un-sequenced features extracted from run 1 and 2. Run 5 (INL-NL): directed LC-MS/MS of unassigned neutral loss showing precursors

**) Numbers calculated from Bioworks 3.2 search results using an in-house software tool (25)

Table 3: Identified phosphorylation sites involved in the Wnt-signaling pathway

Accession number*	Gene name	LC-MS/MS run detected**	Precursor mass (Da)	Charge state	Peptide	Propability***	Sf-score***	Xcorr***	delta CN***
FBgn0001139	gro	1 (DDA 2+)	1182.51506	2	R.NSV S *PADREK.Y	1.83E-03	0.85	3.26	0.31
FBgn0001139	gro	1 (DDA 2+)	1356.67538	2	R.TR S *PLDIENDSK.R	1.52E-06	0.82	3.28	0.43
FBgn0002783	mor	1 (DDA 2+)	1520.71046	2	K.ANAALQSTAS S *PAPGGK.S	1.03E-03	0.70	3.34	0.05
FBgn0015805	Rpd3	1 (DDA 2+)	2141.75841	2	R.IVPENEY S *D S *EDEGEGGR.R	2.53E-05	0.95	6.44	0.09
FBgn0015805	Rpd3	1 (DDA 2+)	2297.85952	2	R.IVPENEY S *D S *EDEGEGGRR.D	2.41E-04	0.56	4.65	0.14
FBgn0022131	aPKC	2 (DDA >2+)	2845.32267	3	K.EGIRPGDTT S *TFCGTPNYIAPEILR.G	1.20E-03	0.53	4.04	0.30
FBgn0003371	sgg	2 (DDA >2+)	1852.84120	3	K.QLLHGEPNVS Y *ICSR.Y	2.23E-03	0.84	4.09	0.08
FBgn0003371	sgg	3 (INL-1)	1020.40338	2	R.TSS S *FAEGNK.Q	8.72E-03	0.55	1.92	0.29
FBgn0032157	CG5899	3 (INL-1)	3177.56749	3	K.LLTALGLDKDQEEQLNNSLNNSIAS S *PAK	2.19E-05	0.93	6.92	0.17
FBgn0032157	CG5899	4 (INL-2)	1981.62937	2	K.DQVYD S *DD S *DSEMSTK.M	2.29E-04	0.95	5.60	0.09
FBgn0015589	Apc	4 (INL-2)	2738.39852	3	R.RNS S *VAGSGQNVDSPVVIPASLQPLR.S	2.30E-04	0.53	3.82	0.09
FBgn0027492	wdb	5 (INL-NL)	2007.98993	2	R.YRNS S *QDVELQQLPPLK.A	4.46E-03	0.90	4.45	0.15

*) Accession number as indicated in Flybase 4.3

**) Run 1 (DDA 2+) and 2 (DDA >2+): only doubly or higher charges peptide ions, respectively, were allowed to trigger CID by DDA. Run 3 (INL-1) and 4 (INL-2): directed LC-MS/MS of un-sequenced features extracted from run 1 and 2. Run 5 (INL-NL): directed LC-MS/MS of unassigned neutral loss showing precursors

***) Search result values obtained from Bioworks 3.2

Figure 1:

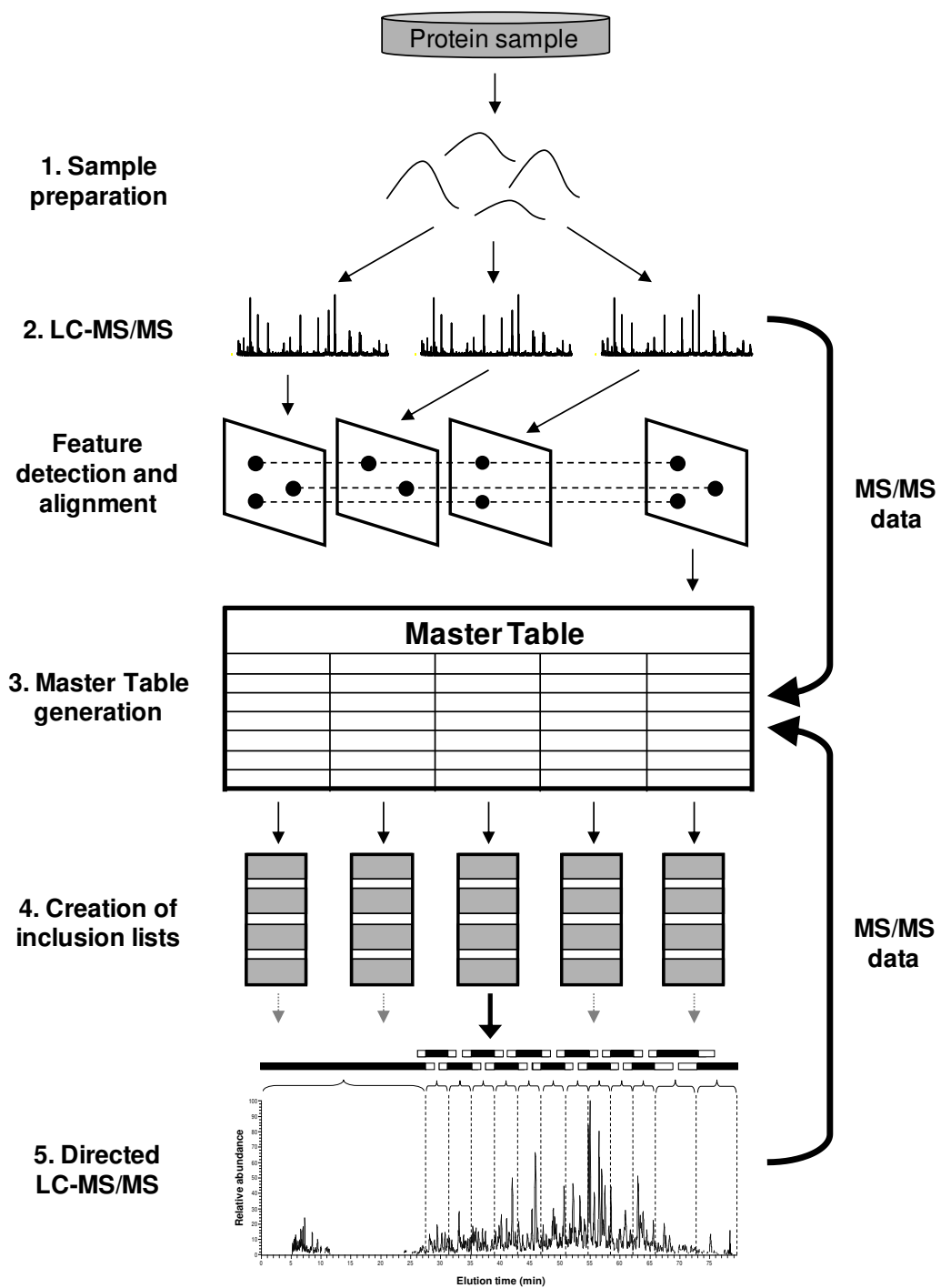


Figure 2:

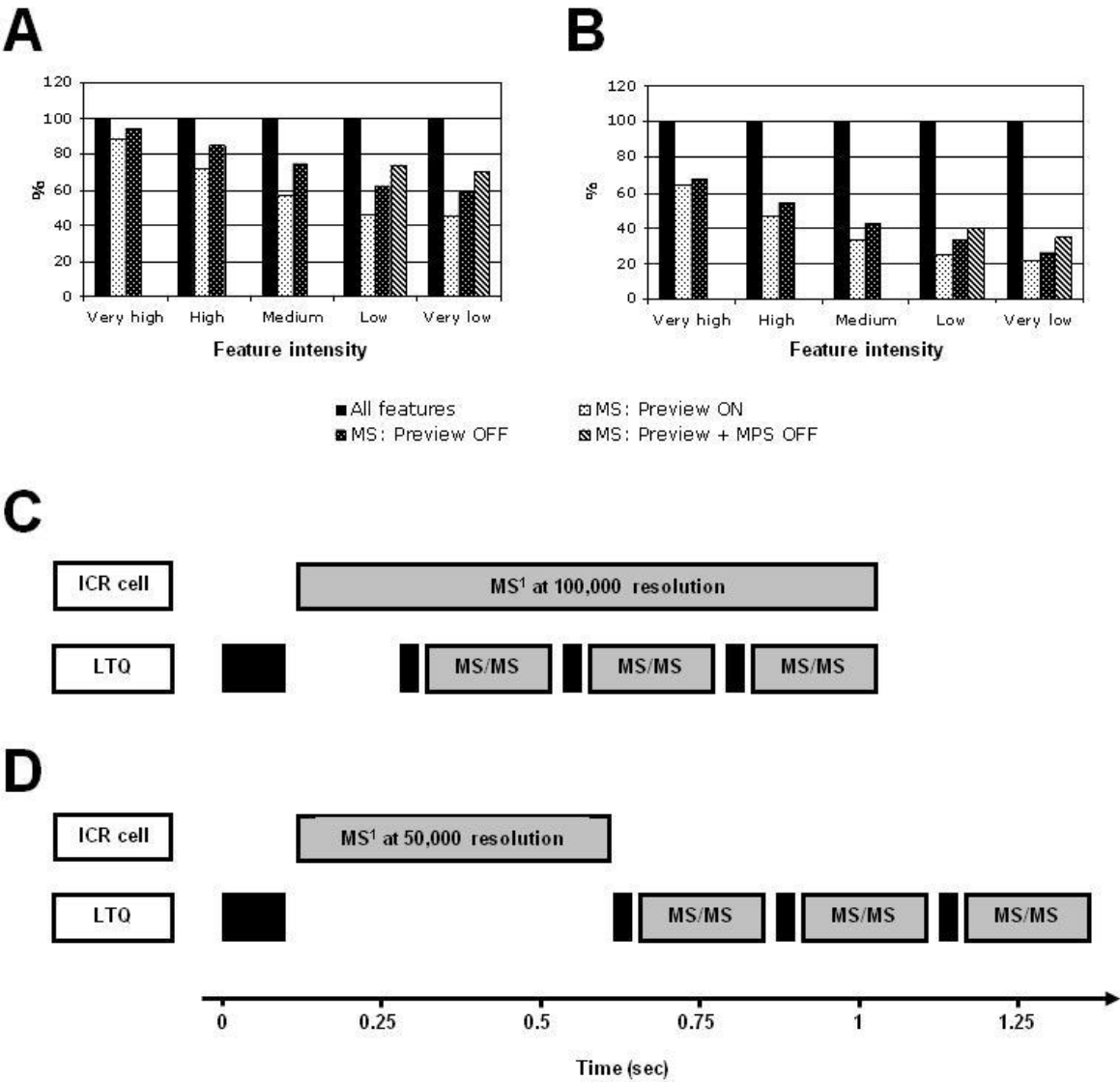


Figure 3:

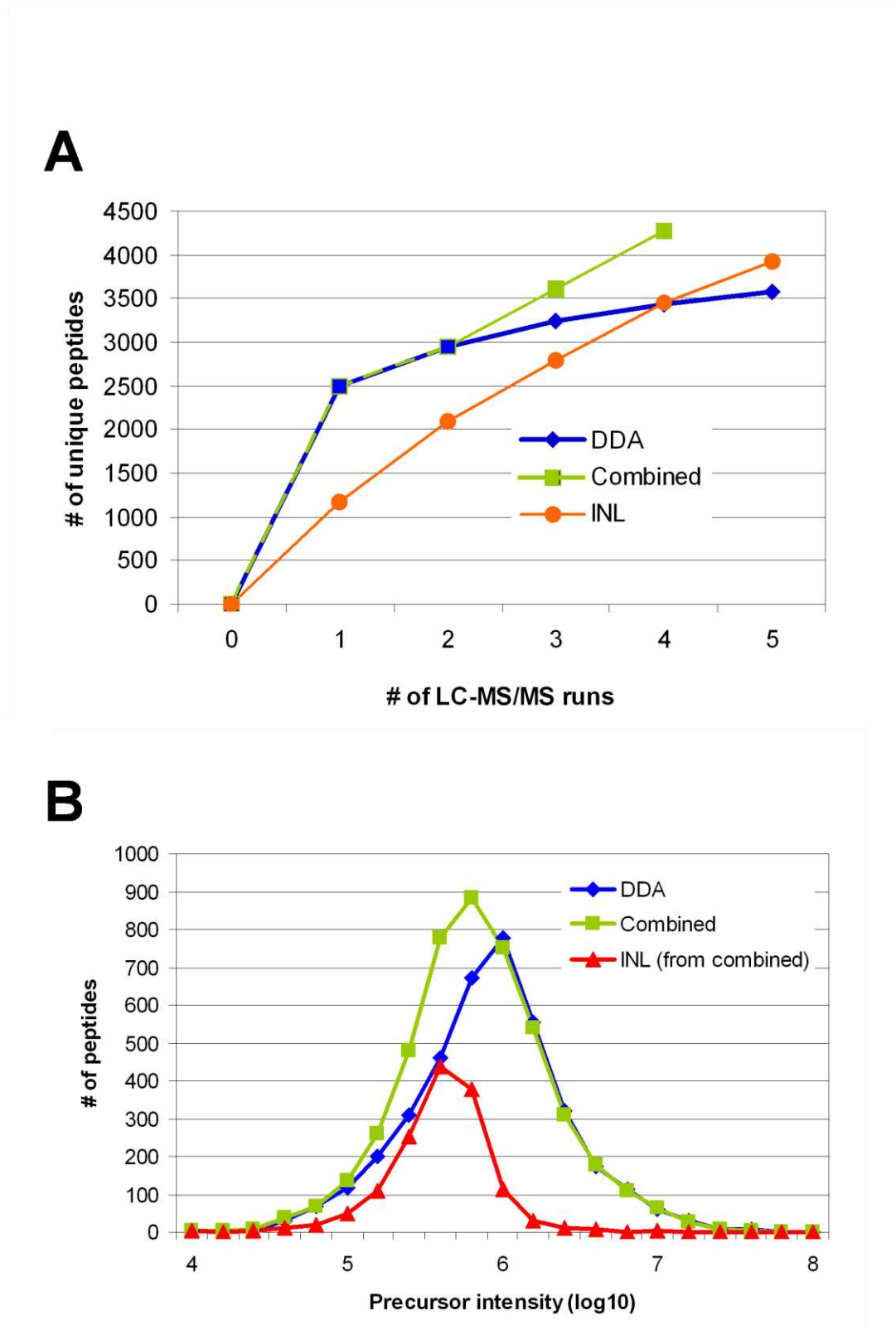
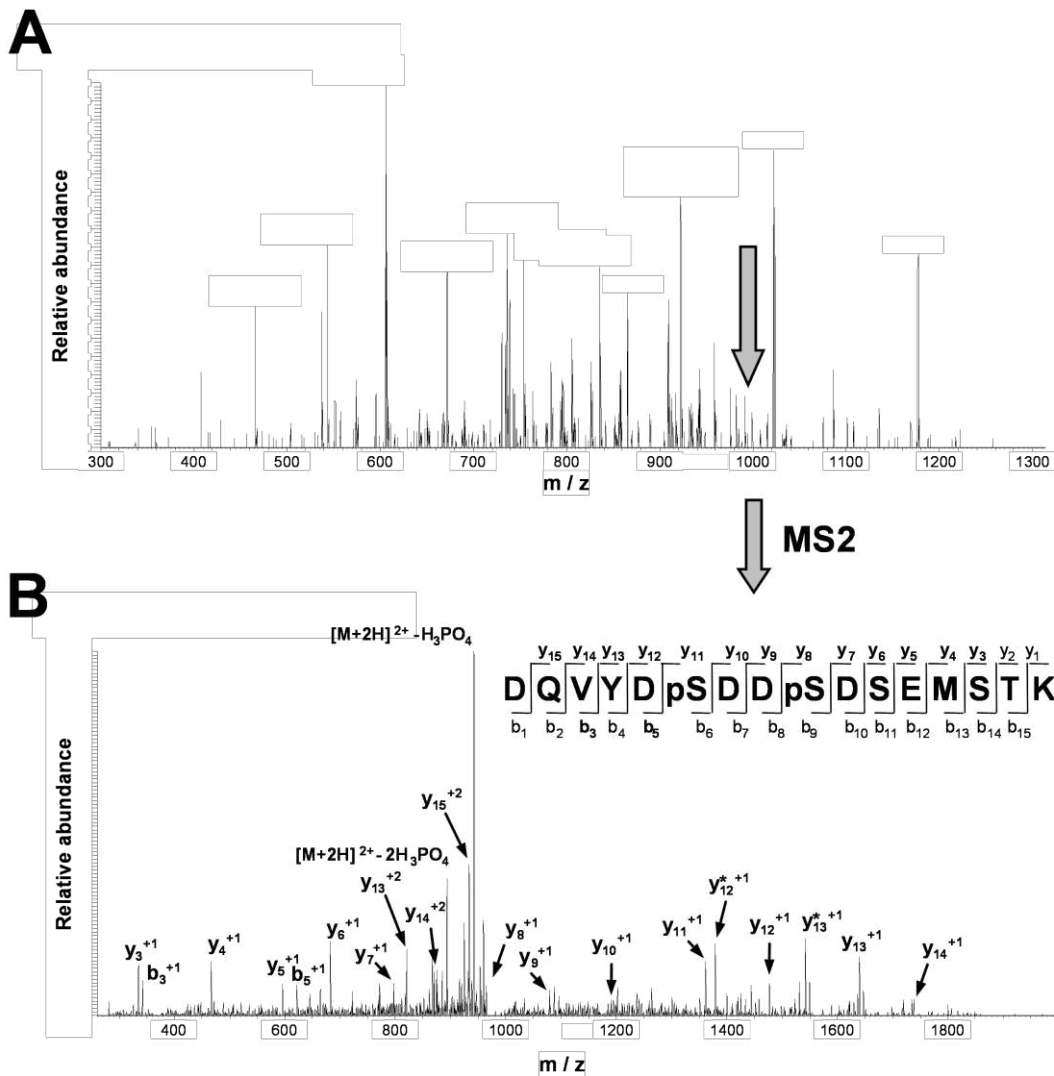


Figure 4:



References

1. Aebersold, R., and Mann, M. (2003) Mass spectrometry-based proteomics. *Nature* 422, 198-207.
2. Gygi, S. P., Rist, B., Gerber, S. A., Turecek, F., Gelb, M. H., and Aebersold, R. (1999) Quantitative analysis of complex protein mixtures using isotope-coded affinity tags. *Nat Biotechnol* 17, 994-999.
3. Ong, S. E., Kratchmarova, I., and Mann, M. (2003) Properties of ¹³C-substituted arginine in stable isotope labeling by amino acids in cell culture (SILAC). *J Proteome Res* 2, 173-181.
4. Schmidt, A., Kellermann, J., and Lottspeich, F. (2005) A novel strategy for quantitative proteomics using isotope-coded protein labels. *Proteomics* 5, 4-15.
5. Brunner, E., Ahrens, C. H., Mohanty, S., Baetschmann, H., Loevenich, S., Potthast, F., Deutsch, E. W., Panse, C., de Lichtenberg, U., Rinner, O., Lee, H., Pedrioli, P. G., Malmstrom, J., Koehler, K., Schrimpf, S., Krijgsveld, J., Kregenow, F., Heck, A. J., Hafen, E., Schlapbach, R., and Aebersold, R. (2007) A high-quality catalog of the *Drosophila melanogaster* proteome. *Nat Biotechnol* 25, 576-583.
6. Kislinger, T., Cox, B., Kannan, A., Chung, C., Hu, P., Ignatchenko, A., Scott, M. S., Gramolini, A. O., Morris, Q., Hallett, M. T., Rossant, J., Hughes, T. R., Frey, B., and Emili, A. (2006) Global survey of organ and organelle protein expression in mouse: combined proteomic and transcriptomic profiling. *Cell* 125, 173-186.
7. Adachi, J., Kumar, C., Zhang, Y., Olsen, J. V., and Mann, M. (2006) The human urinary proteome contains more than 1500 proteins, including a large proportion of membrane proteins. *Genome Biol* 7, R80.
8. de Godoy, L. M., Olsen, J. V., de Souza, G. A., Li, G., Mortensen, P., and Mann, M. (2006) Status of complete proteome analysis by mass spectrometry: SILAC labeled yeast as a model system. *Genome Biol* 7, R50.
9. Kuster, B., Schirle, M., Mallick, P., and Aebersold, R. (2005) Scoring proteomes with proteotypic peptide probes. *Nat Rev Mol Cell Biol* 6, 577-583.
10. Kristensen, D. B., Brond, J. C., Nielsen, P. A., Andersen, J. R., Sorensen, O. T., Jorgensen, V., Budin, K., Matthiesen, J., Veno, P., Jespersen, H. M., Ahrens, C. H., Schandorff, S., Ruhoff, P. T., Wisniewski, J. R., Bennett, K. L., and Podtelejnikov, A. V. (2004) Experimental Peptide Identification Repository (EPIR): an integrated peptide-centric platform for validation and mining of tandem mass spectrometry data. *Mol Cell Proteomics* 3, 1023-1038.
11. Liu, H., Sadygov, R. G., and Yates, J. R., 3rd (2004) A model for random sampling and estimation of relative protein abundance in shotgun proteomics. *Anal Chem* 76, 4193-4201.
12. Domon, B., and Aebersold, R. (2006) Mass spectrometry and protein analysis. *Science* 312, 212-217.
13. Rinner, O., Mueller, L. N., Hubalek, M., Muller, M., Gstaiger, M., and Aebersold, R. (2007) An integrated mass spectrometric and computational framework for the analysis of protein interaction networks. *Nat Biotechnol* 25, 345-352.
14. Domon, B., and Broder, S. (2004) Implications of new proteomics strategies for biology and medicine. *J Proteome Res* 3, 253-260.

15. Bisle, B., Schmidt, A., Scheibe, B., Klein, C., Tebbe, A., Kellermann, J., Siedler, F., Pfeiffer, F., Lottspeich, F., and Oesterhelt, D. (2006) Quantitative profiling of the membrane proteome in a halophilic archaeon. *Mol Cell Proteomics* 5, 1543-1558.
16. Griffin, T. J., Lock, C. M., Li, X. J., Patel, A., Chervetsova, I., Lee, H., Wright, M. E., Ranish, J. A., Chen, S. S., and Aebersold, R. (2003) Abundance ratio-dependent proteomic analysis by mass spectrometry. *Anal Chem* 75, 867-874.
17. Calvo, S., Jain, M., Xie, X., Sheth, S. A., Chang, B., Goldberger, O. A., Spinazzola, A., Zeviani, M., Carr, S. A., and Mootha, V. K. (2006) Systematic identification of human mitochondrial disease genes through integrative genomics. *Nat Genet* 38, 576-582.
18. Picotti, P., Aebersold, R., and Domon, B. (2007) The Implications of Proteolytic Background for Shotgun Proteomics.
19. Gehlenborg, N., Yan, W., Yoo, H., Lee, I., Nieselt, K., Hwang, D., Aebersold, R., and Hood, L. (Manuscript in preparation) Prequips - An Extensible Software Platform for Integration, Visualization and Analysis of LC-MS/MS Proteomics Data.
20. Bodenmiller, B., Mueller, L. N., Mueller, M., Domon, B., and Aebersold, R. (2007) Reproducible isolation of distinct, overlapping segments of the phosphoproteome. *Nat Methods* 4, 231-237.
21. Pedrioli, P. G., Eng, J. K., Hubley, R., Vogelzang, M., Deutsch, E. W., Raught, B., Pratt, B., Nilsson, E., Angeletti, R. H., Apweiler, R., Cheung, K., Costello, C. E., Hermjakob, H., Huang, S., Julian, R. K., Kapp, E., McComb, M. E., Oliver, S. G., Omenn, G., Paton, N. W., Simpson, R., Smith, R., Taylor, C. F., Zhu, W., and Aebersold, R. (2004) A common open representation of mass spectrometry data and its application to proteomics research. *Nat Biotechnol* 22, 1459-1466.
22. Mueller, L. N., Rinner, O., Schmidt, A., Letarte, S., Bodenmiller, B., Brusniak, M. Y., Vitek, O., Aebersold, R., and Muller, M. (2007) SuperHirn - a novel tool for high resolution LC-MS-based peptide/protein profiling. *Proteomics* 7, 3470-3480.
23. Elias, J. E., and Gygi, S. P. (2007) Target-decoy search strategy for increased confidence in large-scale protein identifications by mass spectrometry. *Nat Methods* 4, 207-214.
24. Nesvizhskii, A. I., Keller, A., Kolker, E., and Aebersold, R. (2003) A statistical model for identifying proteins by tandem mass spectrometry. *Anal Chem* 75, 4646-4658.
25. Bodenmiller, B., Mueller, L. N., Pedrioli, P. G., Pflieger, D., Junger, M. A., Eng, J. K., Aebersold, R., and Tao, W. A. (2007) An integrated chemical, mass spectrometric and computational strategy for (quantitative) phosphoproteomics: application to *Drosophila melanogaster* Kc167 cells. *Mol Biosyst* 3, 275-286.
26. Williams, D. K., Jr., and Muddiman, D. C. (2007) Parts-per-billion mass measurement accuracy achieved through the combination of multiple linear regression and automatic gain control in a Fourier transform ion cyclotron resonance mass spectrometer. *Anal Chem* 79, 5058-5063.
27. Masselon, C., Tolmachev, A. V., Anderson, G. A., Harkewicz, R., and Smith, R. D. (2002) Mass measurement errors caused by "local" frequency

- perturbations in FTICR mass spectrometry. *J Am Soc Mass Spectrom* 13, 99-106.
28. Haas, W., Faherty, B. K., Gerber, S. A., Elias, J. E., Beausoleil, S. A., Bakalarski, C. E., Li, X., Villen, J., and Gygi, S. P. (2006) Optimization and use of peptide mass measurement accuracy in shotgun proteomics. *Mol Cell Proteomics* 5, 1326-1337.
 29. Lam, H., Deutsch, E. W., Eddes, J. S., Eng, J. K., King, N., Stein, S. E., and Aebersold, R. (2007) Development and validation of a spectral library searching method for peptide identification from MS/MS. *Proteomics* 7, 655-667.
 30. Larsen, M. R., Thingholm, T. E., Jensen, O. N., Roepstorff, P., and Jorgensen, T. J. (2005) Highly selective enrichment of phosphorylated peptides from peptide mixtures using titanium dioxide microcolumns. *Mol Cell Proteomics* 4, 873-886.
 31. Schroeder, M. J., Shabanowitz, J., Schwartz, J. C., Hunt, D. F., and Coon, J. J. (2004) A neutral loss activation method for improved phosphopeptide sequence analysis by quadrupole ion trap mass spectrometry. *Anal Chem* 76, 3590-3598.
 32. Li, X., Gerber, S. A., Rudner, A. D., Beausoleil, S. A., Haas, W., Villen, J., Elias, J. E., and Gygi, S. P. (2007) Large-scale phosphorylation analysis of alpha-factor-arrested *Saccharomyces cerevisiae*. *J Proteome Res* 6, 1190-1197.
 33. Olsen, J. V., Blagoev, B., Gnäd, F., Macek, B., Kumar, C., Mortensen, P., and Mann, M. (2006) Global, in vivo, and site-specific phosphorylation dynamics in signaling networks. *Cell* 127, 635-648.
 34. Reya, T., and Clevers, H. (2005) Wnt signalling in stem cells and cancer. *Nature* 434, 843-850.
 35. Segditsas, S., and Tomlinson, I. (2006) Colorectal cancer and genetic alterations in the Wnt pathway. *Oncogene* 25, 7531-7537.
 36. Thomas, P. D., Campbell, M. J., Kejariwal, A., Mi, H., Karlak, B., Daverman, R., Diemer, K., Muruganujan, A., and Narechania, A. (2003) PANTHER: a library of protein families and subfamilies indexed by function. *Genome Res* 13, 2129-2141.
 37. Keller, A., Nesvizhskii, A. I., Kolker, E., and Aebersold, R. (2002) Empirical statistical model to estimate the accuracy of peptide identifications made by MS/MS and database search. *Anal Chem* 74, 5383-5392.
 38. Old, W. M., Meyer-Arendt, K., Aveline-Wolf, L., Pierce, K. G., Mendoza, A., Sevinsky, J. R., Resing, K. A., and Ahn, N. G. (2005) Comparison of label-free methods for quantifying human proteins by shotgun proteomics. *Mol Cell Proteomics* 4, 1487-1502.
 39. Goodlett, D. R., Bruce, J. E., Anderson, G. A., Rist, B., Pasa-Tolic, L., Fiehn, O., Smith, R. D., and Aebersold, R. (2000) Protein identification with a single accurate mass of a cysteine-containing peptide and constrained database searching. *Anal Chem* 72, 1112-1118.
 40. Rinner, O., Seebacher, J., Walzthoeni, T., Mueller, L., Beck, M., Schmidt, A., Mueller, M., and Aebersold, R. (2008) Identification of cross-linked peptides from large sequence databases. *Nat Methods* 5, 315-318.
 41. Yi, E. C., Marelli, M., Lee, H., Purvine, S. O., Aebersold, R., Aitchison, J. D., and Goodlett, D. R. (2002) Approaching complete peroxisome characterization by gas-phase fractionation. *Electrophoresis* 23, 3205-3216.